ATTY DOCKET APPLICANT(S)

: RM.WSL

: Le Yi Wang; Hong Wang; and Gang George Yin

SERIAL NO. FILED

: 10/561,074

: May 22, 2006

Art Unit: 4185 Conf. No.: 1880

Examiner: Atia K. Syed

In the Drawing:

Please amend the drawing figures in accordance Annexure 3, attached hereto, which

constitutes sketches presented on twelve (12) replacement drawing sheets annexed hereto.

REMARKS

Amendments are presented herein to improve the form of the subject application and in

response to the Examiner's comments in the above-identified Office Action.

Drawings

Figures 1, 11, 12, 14, 15, 19, 20 and 22-26 are considered by the Examiner to be

objectionable because the size of the figures and the font size of the legends are too small to

properly and clearly review the figures and zooming into the figure is not possible due to poor

resolution. The Examiner has required that Applicants provide corrected drawing sheets in

compliance with 37 C.F.R. § 1.121(d) in reply to the Office action to avoid abandonment of the

application.

The Examiner advises that any amended replacement drawing sheet should include all

of the figures appearing on the immediate prior version of the sheet, even if only one figure is

being amended. The figure or figure number of an amended drawing should not be labeled as

"amended." If a drawing figure is to be canceled, the appropriate figure must be removed from

the replacement sheet, and where necessary, the remaining figures must be renumbered and

appropriate changes made to the brief description of the several views of the drawings for

consistency. Additional replacement sheets may be necessary to show the renumbering of the

remaining figures. According to the Examiner, each drawing sheet submitted after the filing date

of an application must be labeled in the top margin as either "Replacement Sheet" or "New

Sheet" pursuant to 37 C.F.R. § 1.121(d). If the changes are not accepted by the Examiner,

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Applicants will be notified and informed of any required corrective action in the next Office action. The objection to the drawings will not be held in abeyance by the Examiner.

APPLICANTS' RESPONSE

Applicants have amended the drawing figures to conform to the requirements stated by the Examiner. More specifically, the drawings have been redrawn in a larger size and all text has been replaced with higher resolution text. No new matter has been added to the drawing figures, and all REPLACEMENT SHEETS are attached and have been identified accordingly, as required by the Examiner.

The resulting distribution of the drawing figures is as follows:

- Fig. 1 -Formerly on same sheet with Figs. 2a and 2b. Now, enlarged and clarified Fig. 1 is on a separate REPLACEMENT SHEET, and Figs, 2a and 2b are together on a separate REPLACEMENT SHEET;
- Formerly on same sheet with Fig. 12. Now, enlarged and clarified Fig. Fig. 11 -11 is on a separate REPLACEMENT SHEET;
- Fig. 12 -Formerly on same sheet with Fig. 11. Now, enlarged and clarified Fig. 12 is on a separate REPLACEMENT SHEET;
- Fig. 14 -Formerly on same sheet with Fig. 15. Now, enlarged and clarified Fig. 14 is on a separate REPLACEMENT SHEET;
- Fig. 15 -Formerly on same sheet with Fig. 14. Now, enlarged and clarified Fig. 15 is on a separate REPLACEMENT SHEET;
- Formerly on same sheet with Fig. 20. Now, enlarged and clarified Fig. Fig. 19 -19 is on a separate REPLACEMENT SHEET;
- Fig. 20 -Formerly on same sheet with Fig. 19. Now, enlarged and clarified Fig. 20 is on a separate REPLACEMENT SHEET;

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Fig. 22 - Formerly on same sheet with Figs. 23 and 24. Now, enlarged and

clarified Fig. 22 is on a separate REPLACEMENT SHEET with Fig. 23;

Fig. 23 - Formerly on same sheet with Figs. 22 and 24. Now, enlarged and

clarified Fig. 23 is on a separate REPLACEMENT SHEET with Fig. 22;

Fig. 24 - Formerly on same sheet with Figs. 22 and 23. Now, enlarged and

clarified Fig. 24 is on a separate REPLACEMENT SHEET;

Fig. 25 - Formerly on same sheet with Fig. 26. Now, enlarged and clarified Fig.

25 is on a separate REPLACEMENT SHEET; and

Fig. 26 - Formerly on same sheet with Fig. 25. Now, enlarged and clarified Fig.

26 is on a separate REPLACEMENT SHEET.

In view of the foregoing, it is respectfully asserted that the Examiner's objections to the drawings have been overcome.

Claim Rejections - 35 U.S.C. § 101

The Examiner states that "claims 1-10 stand rejected under 35 U.S.C. § 101 because the claimed invention is considered by the Examiner to be directed to non-statutory subject matter."

The Examiner further states that claims 1-6 stand rejected under 35 U.S.C. § 101 because the claimed invention is considered by the Examiner to be directed to non-statutory subject matter. In particular, the Examiner states that "claims 1-9 are drawn to a process." The Examiner continues the comment by noting that under 35 U.S.C. § 101 a process must 1) be tied to another statutory class (such as a particular apparatus) or 2) transform underlying subject matter (such as an article or materials) to a different state or thing. The claimed process steps are not considered by the Examiner to transform underlying subject matter.

The Examiner continues the comment by noting that to qualify as a 35 U.S.C. § 101 statutory process, the claims should positively recite the other statutory class (apparatus or thing) to which it is tied, for example by identifying the apparatus that accomplishes the method steps.

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In this regard, the Examiner directs Applicants to http://www.uspto.gov/web/offices/pac/dapp/-

opla/preognotice/section 101 05 15 2008.pdf.

APPLICANTS' RESPONSE

Applicants respectfully disagree with several aspects of the Examiner's comments. First

it is noted that the Examiner has rejected claims 1-10 under 35 U.S.C. § 101. This rejection

should not be applied to independent claim 7, which is clearly an apparatus claim, and its

associated dependent claims 8-10.

Second, for the reason indicated above, Applicants disagree with the Examiner's

assertion that "claims 1-9 are drawn to a process." According to Applicants' records, claims 7-

10 are apparatus claims, not process claims. More specifically, independent claim 7 specifies

a first memory; a second memory; a third memory; a signal combiner arrangement; a limiter

; and a virtual anesthesia monitor. None of these claimed elements constitutes a process step.

With respect to independent claims 1 and 4, these claims have been amended to conform

with the Examiner's stated requirements imposed by 35 U.S.C. § 101. The claimed process is

now directed to the use of a computing machine having a memory. The process requires the

entry of data corresponding to coefficients C_1 , C_2 , C_3 , as well as time periods τ_n (initial time delay

after drug infusion) and T_p (time constant representing speed of response) to be entered into the

memory of the computing machine. As such, this is a 35 U.S.C. § 101 process that is "... tied

to another statutory class ...," and therefore complies with the discussion on Interim Guidelines

set forth in the Memorandum of May 15, 2008 from John J. Love, Deputy Commissioner for

Patent Examination Policy, to the Technology Center Directors. Specifically, the process of

amended independent claim 1 is tied to the statutory machine class of 35 U.S.C. § 101.

Dependent claims 2, 3, 5 and 6 depend from amended independent claims 1 and 4, and have been

correspondingly similarly amended.

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In view of the foregoing, it is respectfully asserted that the Examiner's rejection of claims

1-6 under 35 U.S.C. § 101 has been overcome.

With respect to independent claim 7, this claim has been amended to specify with greater

precision that the claimed subject matter corresponds to "apparatus," rather than a "system."

Independent claim 4 has been amended to state the statutory machine class of 35 U.S.C.

§ 101 with greater specificity. More specifically, the preamble of this apparatus claim has been

amended to identify the claimed subject matter as "apparatus," rather than a "system."

In view of the foregoing, the Examiner's rejection of claims 1-10 under 35 U.S.C. § 101

is believed to have been overcome.

Claim Rejections - 35 U.S.C. § 112, Second Paragraph

Claims 7-10 stand rejected under 35 U.S.C. § 112, second paragraph, as being indefinite

for failing to point out with particularity and claim distinctly the subject matter that Applicants

regard as the invention. According to the Examiner, the metes and the bounds of the claimed

structure can not be determined by the disclosure.

The Examiner states that Applicants have claimed a signal combiner, a limiter, and the

virtual anesthesia monitor. However, it is unclear to the Examiner if the claimed signal

combiner, limiter and virtual anesthesia monitor are structures, or algorithms, or software (data

structures). If the determination means is an algorithm or software then the term/limitation will

not be given patentable weight because it lacks structure that would be attributed to the apparatus

claims.

The Examiner notes that the word "for" in the claims can may be interpreted as intended

use. Intended use/functional language does not, according to the Examiner, require that a

reference specifically teach the intended use of the element. A recitation of the intended use of

the claimed invention must, according to the Examiner, result in a structural difference between

the claimed invention and the prior art in order to patentably distinguish the claimed invention

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from the prior art. If the prior art structure is capable of performing the intended use then,

according to the Examiner, it meets the claim.

APPLICANTS' RESPONSE

Applicants respectfully disagree with the Examiner's contention that algorithms and data

structures in the present case are not entitled to patentable weight. Moreover, it is respectfully

further asserted that the Examiner incorrectly contends that these elements would "[lack]

structure that would be attributed to ... apparatus claims."

In regard of the foregoing it is noted that independent claim 7 has since the time of filing

of the present case specified its claimed subject matter in the context of a "system," that contains

elements that unquestionably constitute apparatus. These include, for example, the first, second,

and third memories. The remaining elements, specifically the signal combiner arrangement, the

limiter, and the virtual anesthesia monitor, are described in the present application at, for

example, Fig. 1, which shows a plurality of signals being combined in a computer, and in

corresponding portions of the specification (e.g., page 5, lines 5-16). It is evident from the

figures and the specification that these elements, i.e., the signal combiner arrangement, the

limiter, and the virtual anesthesia monitor, as well as the limiter, are implemented in a computer.

It is well-established that a computer is a machine that assumes machine characteristics that are

responsive to the program it is running.

In the present case, the claimed elements identified by the Examiner are not merely

algorithms or data structures, but instead constitute machine elements having every characteristic

of apparatus. Accordingly, for these and other reasons, it is respectfully asserted that the metes

and bounds of these elements can readily be determined from the specification and drawings, and

accordingly, the rejection of claims 7-10 by the Examiner under 35 U.S.C. § 112, second

paragraph, has been traversed.

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The foregoing notwithstanding, Applicants have amended claims 7-10 to specify the

subject matter therein claimed as "apparatus," which clearly is a statutory class, the claimed

combination of elements being entitled to patentable weight.

Claim Rejections - 35 U.S.C. § 102(b)

Claims 1-10 stand rejected under 35 U.S.C. § 102(b) as specifying subject matter

considered by the Examiner to be anticipated by Kangas, et al. (US 5,775,330) herein after

referred as Kangas, et al.

Regarding claims 1-6, the Kangas, et al. reference is considered by the Examiner to

disclose a method of predicting the anesthetic depth of a person by analyzing his EEG signals

(claim 1).

Regarding claims 7-10, the Kangas, et al. reference is considered by the Examiner to

disclose an apparatus that recodes the data (column 7, lines 3-37), process the EEG signals and

transforms them to predict the anesthetic depth of a patient (claim 13). The invention of the

Kangas, et al. reference includes a memory that is used to record data during surgery for seven

different subjects (column 7, lines 3-9).

APPLICANTS' RESPONSE

Briefly, the Kangas, et al. reference describes a method and apparatus for collecting EEG

data, reducing the EEG data into coefficients, and correlating those coefficients with a depth of

unconsciousness or anesthetic depth, and which obtains a bounded first derivative of anesthetic

depth to indicate trends. An artificial neural network based method continuously analyzes EEG

data to discriminate between awake and anesthetized states in an individual and continuously

monitors anesthetic depth trends in real-time. This enables an anesthesiologist to respond to

changes in anesthetic depth of the patient during surgery and to administer the correct amount

of anesthetic. The use of brain wave data processed by a trained neural network ascertains the

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level of consciousness of an individual and a consciousness trend before during and after

exposure to anesthetic.

The foregoing notwithstanding, the system described in the Kangas, et al. reference

cannot determine whether the patient is adequately anesthetized. The adequacy of anesthesia is

a difficult problem for anesthesiologists who occasionally fail despite the fact that they attempt

to over anesthetize the patient to ensure adequate anesthesia.

The Kangas, et al. reference is directed to anesthesia depth measurement and calculation

only. The word "prediction" is used in regard of the known system only in the sense that EEG

signals are "analyzed" to "calculate" anesthesia depth and its derivatives. This known system,

however, does not address the issue of how a drug input strategy will influence the anesthesia

depth in the future. On the contrary, the present claimed invention achieves drug impact

prediction. The system of the Kangas, et al. reference does not possess a "prediction" capability,

as does the present invention, but rather is a method of calculating actual (not predicted)

anesthesia depth. In short, the Kangas, et al. reference teaches subject matter already in use in

FDA approved devices, such as the BIS Monitor by Aspect, and Entropy Monitor by GE, which

are employed for anesthesia depth measurements and which produce a corresponding display to

the operator. In contrast, anesthesia depth measurements are the input signals to the present

claimed system.

In addition to the foregoing, it is noted that drug impact prediction depends on the

response to drugs by each individual patient. Thus, there is a need for individual patient dynamic

models. The system described in the Kangas, et al. reference has nothing to do with patient

models. In fact, it is admitted in the Kangas, et al. reference that the invention therein described

cannot determine whether the patient is adequately anesthetized. It is therefore respectfully

asserted that the Examiner has incorrectly characterized the Kangas, et al. reference as

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"predicting the anesthetic depth of a person by analyzing his EEG signals" or as useful to

"process the EEG signals and transform them to predict the anesthetic depth of a patient."

With reference to amended independent claim 1, it is noted that this claim specifies a

"method of using a computing machine having a memory to assist a human expert in reducing

predictable variations in the depth of anesthesia during the administration of a medical anesthesia

drug to a patient." Nothing in the Kangas, et al. reference achieves, teaches, or suggests such

prediction.

With reference to amended independent claim 4, it is noted that this claim specifies a

"method of using a computing machine having a memory to determine a model that corresponds

to a predicted response of a patient to anesthesia drug delivery." Nothing in the Kangas, et al.

reference teaches or suggests the determination of a model of a patient response, or a prediction

of the patient's response to anesthesia.

With respect to amended independent claim 7, it is noted that this claim specifies

"apparatus for determining a predicted response of a patient to the administration of an

anesthesia drug." Again, nothing in the Kangas, et al. reference teaches or suggests the

predicting of a patient's response to anesthesia.

In view of the foregoing, it is respectfully asserted that the Examiner's rejection of claims

1-10 under 35 U.S.C. § 102(b) has been overcome, and that these claims are in allowable

condition.

Prior Art Cited But Not Applied

The prior art made of record and not relied upon is considered by the Examiner to be

pertinent to Applicants' disclosure. In this regard, the Examiner cited the following references

as disclosing related limitations of the applicant's claimed and disclosed invention.

Nomura; Takashi, et al. (US 5964713 A), is considered by the Examiner to disclose a

method and apparatus to predict the anesthetic depth of a patient undergoing surgery. Briefly,

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this reference discloses a system that achieves only anesthesia depth measurement and

calculation. The word "prediction" is used only in the sense that it can "analyze" body signals

(such as blood pressures, body temperature, and respiratory rates) to "calculate" anesthesia

depth. The known system does not address the issue of how a drug input strategy will influence

the anesthesia depth in the future. Thus, this known arrangement is not capable of generating

a "prediction," but instead is a method of calculating anesthesia depth. In addition, the subject

matter of this reference teaches nothing regarding patient models.

Tang; Sharon S., et al. (US 6658396 B1), is considered by the Examiner to disclose an

apparatus for predicting the optimal dosage for a particular patient while considering a plurality

of factors.

In this known arrangement, a computerized neural network drug dosage estimator

predicts what will happen when a patient, or a patient population, is administered a drug dosage

deviating from predicted optimal dosage. For example, if an individual patient of certain

characteristics does not exhibit desired therapeutic response at a certain (possibly even the

predicted optimal) drug dosage level, then the question arises whether the dosage should be

increased by 10%, or by 25%, or by 50%, or even by 100%? The neural network drug dosage

estimator of the Tang, et al. reference helps to answer this question.

The known arrangement is embodied in a computerized method of predicting an optimal

dosage of a particular drug for a particular patient in consideration of previously determined

optimal dosages of the drug for members of a patient population. More particularly, the known

system requires that a neural network having an architecture of one or more slabs be

programmed. The slabs collectively relate "input data" to "output data". The "input data"

includes at least a selected three (3) of a person's traits drawn from at least two (2) of the three

(3) groups of information, specifically:

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Group 1 overt indications of (1a) age, (1b) gender, (1c) race, (1d) ethnicity, (1e) diet

type, (1f) height, (1g) weight, and (1h) body surface area;

Group 2 medical diagnostic indications of (2a) blood pressure, (2b) use of a drug other

than the particular drug at the same time as use of the particular drug, (2c) fitness,

(2d) peptide levels, and (2e) genetic predisposition to a particular disease; and

Group 3 pharmacological indications of (3a) pharmacokinetic parameters, (3b)

pharmacodynamic parameters.

The number and diversity of traits quantitatively distinguish the system described in the

Tang, et al. reference over systems that employ a family of curves relating recommended dosage

by age, weight and sex (i.e., by three traits). The "output data" is the clinically-determined

optimal drug dosage for the same person.

The system described in the Tang, et al. reference is quite complex, as it uses neural

network models that inevitably require determination of a large number of model parameters.

They require a large and very rich data set for training and tuning. Consequently, they cannot

be established in real-time during surgical procedures. The present invention, on the other hand,

utilizes only a very small number of parameters that are computationally feasible to be

established in real time during anesthesia procedures.

A further advantage of the present invention over that described in the Tang, et al.

reference is the interaction of the model with physicians. More specifically, model parameters

in neural networks do not carry any physiological meaning. Consequently, they cannot be

communicated to physicians. The system of the present invention uses parameters derived from

anesthesiologists' viewpoints of anesthesia depth responses, and hence they have direct and clear

physiological meanings, such as time delay, response speed, drug sensitivity. They allow

physicians to understand the model operation, to set safety bounds of parameters, and to

calibrate. The elements of the claims of the present invention relate directly to such parameters.

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A still further advantage of the present invention over the system described in the Tang,

et al. reference relates to the data sets needed for establishing the models. In the known

arrangement, the models must be learned and tuned by using large population data. As stated

in the Tang, et al. patent "the number of cases required to train the neural network is on the order

of a minimum of 500 to 2,000, and more commonly 10,000." To establish an individual patient's

model, only one patient's data in one anesthesia procedure can be used. More specifically, the

inventive system establishes a patient's unique dynamic model in real-time, without any

requirements from other patients' models.

Yet another advantage of the present invention over the system of the Tang, et al.

reference relates to real-time and individualized patient models. Drug impact prediction depends

on each individual patient's response to drugs, hence requires individual patient's dynamic

models. The system of the Tang, et al. reference uses models that are established by using large

data sets from a large patient population (such as 10000 cases). Hence, they constitute average

population models, not real prediction for an individual but rather guesses of most probable

outcomes based on population studies of large numbers of patients. Large variations in patient

dynamics and drug responses make population-based average models unusable for real-time

anesthesia depth prediction, control, and management. In other words, the system of the Tang,

et al. reference cannot be used to derive an individual patient's model using real-time data in one

anesthesia procedure.

In view of the foregoing, neither the Nomura, et al. nor Tang, et al. references teach or

suggest the claimed invention.

Conclusion

In view of the foregoing, it is respectfully requested that the Examiner reconsider the

present application, allow the claims, and pass the application for issue. If the Examiner believes

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that the prosecution of this case can be expedited by a telephone interview, the Examiner is requested to call attorney for Applicant(s) at the telephone number indicated hereinbelow.

Respectfully submitted,

Report A Monroto

Raphael A. Monsanto Reg. No. 28,448 Rohm & Monsanto, P.L.C. 12 Rathbone Place Grosse Pointe, MI 48230 Telephone (313) 884-4805 Telecopier (313) 884-4806

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enc Annexure 1 - Claims Rewritten to Show Amendments

Annexure 2 - Specification Rewritten to Show Amendments

Annexure 3 - Replacement Sheets of Drawing Figures (12 sheets)

File: ROA-d01.WSL